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July 14, 2000

Case Docket No. C150.12.3B



Assistant Commissioner for Patents  
Washington, D.C. 20231

Transmitted herewith for filing is the patent application of:

Inventor: Peter Nash, John W. Rosevear and Donald L. Robinson

For : IMMUNOGEN ADHERENCE INHIBITOR AND METHOD  
OF MAKING AND USING SAME

Enclosed are:

  x     0   sheets of drawing(s) (informal)

  x    32  pages of specifications

  x   Abstract of the Disclosure.

  x   Declaration

  x   A verified independent inventor and small business establishment statement to establish small entity status under 37 CFR §1.9 and 37 CFR §1.27.

       Disclosure statement with        patent copies.

The filing fees have been calculated: \$ 345 (Small entity \$345 - Large entity \$690)


	(Col. 1)	(Col. 2)	SMALL ENTITY	LARGE ENTITY	
TOTAL CLAIMS	23	- 20	= * 3	x 9 = \$ <u>27</u>	x 18 = \$ <u>-</u>
INDEP. CLAIMS	3	- 3	= * 1	x 39 = \$ <u>6</u>	x 78 = \$ <u>-</u>

\* If the difference in Col. 1 is less than zero, enter "0" in Col. 2

Enclosed is a check in the amount of \$ 345 to cover the filing fee of this application.

The Commissioner is hereby authorized to change any additional filing fees required under 37 CFR §1.16, any patent application processing fees required under 37 CFR §1.17 or credit any overpayment to Deposit Account No. 02-3510. A duplicate copy of this sheet is enclosed.

BURD, BARTZ & GUTENKAUF

  
Robert W. Gutenkauf Reg. No. 25,681  
1300 Foshay Tower  
Minneapolis, MN 55402  
(612) 332-6581

09/616843 - 07/14/00

**VERIFIED STATEMENT CLAIMING SMALL ENTITY STATUS  
(37 CFR 1.9(f) & 1.27(b))--INDEPENDENT INVENTOR**

Docket Number (Optional)

C150.12.3B

Applicant or Patentee: Peter Nash et al.

Serial or Patent No.: \_\_\_\_\_

Filed or Issued: \_\_\_\_\_

Title: IMMUNOGEN ADHERENCE INHIBITOR AND METHOD OF MAKING AND USING SAME

As a below named inventor, I hereby declare that I qualify as an independent inventor as defined in 37 CFR 1.9(c) for purposes of paying reduced fees to the Patent and Trademark Office described in:

- ☒ the specification filed herewith with title as listed above.
- ☐ the application identified above.
- ☐ the patent identified above.

I have not assigned, granted, conveyed or licensed and am under no obligation under contract or law to assign, grant, convey or license, any rights in the invention to any person who would not qualify as an independent inventor under 37 CFR 1.9(c) if that person had made the invention, or to any concern which would not qualify as a small business concern under 37 CFR 1.9(d) or a nonprofit organization under 37 CFR 1.9(e).

Each person, concern or organization to which I have assigned, granted, conveyed, or licensed or am under an obligation under contract or law to assign, grant, convey, or license any rights in the invention is listed below:

- ☐ No such person, concern, or organization exists.
- ☒ Each such person, concern or organization is listed below.

Camas, Incorporated  
1313 Fifth Street SE, Suite 219  
Minneapolis, MN 55414

Separate verified statements are required from each named person, concern or organization having rights to the invention availing to their status as small entities. (37 CFR 1.27)

I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. (37 CFR 1.28(b))

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.

John W. Rosevear, deceased

Peter Nash

by his personal representative

Donald L. Robinson

NAME OF INVENTOR

NAME OF INVENTOR

NAME OF INVENTOR

Peter Nash  
Signature of inventor

Donald L. Robinson  
Signature of inventor

Donald L. Robinson  
Signature of inventor

July 13, 2000  
Date

July 13, 2000  
Date

July 13, 2000  
Date

**VERIFIED STATEMENT CLAIMING SMALL ENTITY STATUS**  
**(37 CFR 1.9(f) & 1.27(c))--SMALL BUSINESS CONCERN**

Docket Number (Optional)  
C150.12.3B
Applicant or Patentee: Peter Nash et al.

Serial or Patent No.: \_\_\_\_\_

Filed or Issued: \_\_\_\_\_

Title: IMMUNOGEN ADHERENCE INHIBITOR AND METHOD OF MAKING AND USING SAME

I hereby declare that I am

☐ the owner of the small business concern identified below:☒ an official of the small business concern empowered to act on behalf of the concern identified below:NAME OF SMALL BUSINESS CONCERN Camas, IncorporatedADDRESS OF SMALL BUSINESS CONCERN 1313 Fifth Street, SE, Suite 219  
Minneapolis, MN 55414

I hereby declare that the above identified small business concern qualifies as a small business concern as defined in 37 CFR 1.21.2, and reproduced in 37 CFR 1.9(d), for purposes of paying reduced fees to the United States Patent and Trademark Office, in that the number of employees of the concern, including those of its affiliates, does not exceed 500 persons. For purposes of this statement, (1) the number of employees of the business concern is the average over the previous fiscal year of the concern of the persons employed on a full-time, part-time or temporary basis during each of the pay periods of the fiscal year, and (2) concerns are affiliates of each other when either, directly or indirectly, one concern controls or has the power to control the other, or a third party or parties controls or has the power to control both.

I hereby declare that rights under contract or law have been conveyed to and remain with the small business concern identified above with regard to the invention described in:

☒ the specification filed herewith with title as listed above.☐ the application identified above.☐ the patent identified above.

If the rights held by the above identified small business concern are not exclusive, each individual, concern or organization having rights in the invention must file separate verified statements averring to their status as small entities, and no rights to the invention are held by any person, other than the inventor, who would not qualify as an independent inventor under 37 CFR 1.9(c) if that person made the invention, or by any concern which would not qualify as a small business concern under 37 CFR 1.9(d), or a nonprofit organization under 37 CFR 1.9(e).

Each person, concern or organization having any rights in the invention is listed below:

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Separate verified statements are required from each named person, concern or organization having rights to the invention averring to their status as small entities. (37 CFR 1.27)

I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. (37 CFR 1.28(b))

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.

NAME OF PERSON SIGNING Peter NashTITLE OF PERSON IF OTHER THAN OWNER PresidentADDRESS OF PERSON SIGNING 1313 Fifth St. SE, Minneapolis, MN 55414SIGNATURE Peter Nash DATE 7/13/00

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**UNITED STATES UTILITY PATENT APPLICATION**  
**OF**

**Peter (nmi) Nash,**

**John W. Rosevear, deceased,**  
**by Donald L. Robinson, his legal representative, and**

**Donald L. Robinson**

**Residents, respectively, of Eden Prairie, Edina and Bloomington, Minnesota**

**Citizens of the United States of America**

**Title: IMMUNOGEN ADHERENCE INHIBITOR**  
**AND METHOD OF MAKING AND**  
**USING SAME**

**Reference is made to Provisional Patent Applications:**  
**Serial No. 60/143,985 filed July 15, 1999**  
**Serial No. 60/201,268 filed May 2, 2000**

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## FIELD OF THE INVENTION

## BACKGROUND OF THE INVENTION

This invention is directed to microbial adherence inhibitor, in the form of fowl egg antibodies, for substantially preventing the attachment or adherence of colony-forming immunogens or haptens in the rumen and intestinal tract of host food animals, to the method of producing such adherence inhibitors, and to the methods of using such inhibitors to: (1) promote the growth of food animals by improving feed conversion rates by decreasing the waste of dietary protein caused by the presence of certain colony-forming protein-wasting organisms in food animals, and (2) to substantially reduce or eliminate the incidence of illnesses caused by the presence of certain illness-causing colony-forming immunogens or haptens in meat from food animals, which are not themselves subject to the targeted illness, and in other food stuffs.

Common bacterial immunogens which cause dramatic decreases in an animal's ability to utilize dietary protein include but are not limited to Peptostreptococcus anaerobius, Clostridium aminophilum, and Clostridium sticklandii. According to Russell (USDA-ARS, May 1993) these organisms, and others disclosed therein, have been collectively responsible for wasting up to 25 percent of the protein in cattle diets. This is a loss of as much as \$25 billion annually to cattle producers and is especially apparent in "grazing animals which are often deficient in protein, even though their protein intake appears to be adequate". As the host consumes protein in the diet, these deleterious organisms wastefully degrade the protein to ammonia which is converted to urea by the liver and kidneys and thus lost to the host when excreted as urine. These deleterious organisms also compete with beneficial organisms which the host needs for the efficient utilization of ammonia. In addition, they need other beneficial organisms in the rumen

the greater the ammonia utilization.

The principal objective of the present invention is to substantially prevent the colonization of deleterious organisms such as P. anaerobius, C. sticklandii and C. aminophilum as well as the growth of such organisms in the rumen and the intestinal tracts of food animals resulting in their substantial elimination from the animal by the administration of the fowl egg antibody to the specific organisms.

Common bacterial immunogens which cause food borne illness in humans include E. coli, Listeria, Salmonella and Campylobacter, all of which produce flu-like symptoms such as nausea, vomiting, diarrhea and/or fever, and in some cases kidney damage or death. In recent years foodstuffs contaminated with these bacteria have caused gastro-intestinal distress in tens or hundreds of thousands of people and the recall and destruction of millions of pounds of food. The resulting economic loss has been staggering. Especially daunting as a public health threat has been E. coli 0157:H7, a pathogenic strain of the common gut bacterium, first identified in 1982. The bacteria are carried in the intestinal tracts of food animals and expelled in their feces. From there, the bacteria enter the food supply, not only in the meat of those animals, but foods such as milk, fruit juices, lettuce, alfalfa sprouts, radishes, and others.

Haptens are partial or incomplete immunogens such as certain toxins, which cannot by themselves cause antibody formation but are capable of combining with specific antibodies. Such haptens may include bacterial toxin, yeast mold toxin, viruses, parasite toxins, algae toxins, etc.

Other colony-forming organisms include Actinomycetes, Streptococcus, Bacteriodes such as B. ruminicola, Cryptococcus and yeasts and mold.

Another principal object of the present invention is to substantially prevent the adherence

of immunogens, such as E. coli 0157:H7, or haptens, and the colonization and growth of such immunogens or haptens in the rumen or intestinal tracts of food animals, and substantial elimination of the immunogen or hapten from the feces of the animals, by the administration to the animals of fowl egg antibody to the specific immunogen or hapten.

## **THE PRIOR ART**

The production of avian egg antibody for the diagnosis or treatment of specific conditions has been known. The production of avian egg antibody for the inhibition of organisms, specifically the colonization of non-illness- causing protein-wasting organisms, and the adherence and colonization of illness-causing immunogens is not suggested.

Representative prior art patents include the following:

Polson, U.S. Patent 4,550,019

Stolle, et al U.S. Patent 4,748,018

Tokoro, U.S. Patent 5,080,895

Carroll, U.S. Patent 5,196,193

Lee, U.S. Patent 5,367,054

Coleman, U.S. Patent 5,585,098

Stolle, et al U.S. Patent 5,753,268

Raun, U.S. Patent 3,794,732, discusses the uses of polyester antibiotics in ruminant rations to improve the utilization of feed in ruminant animals. This specifically addresses the use of antibiotics in ruminant animals as growth promotants.

Raun, U.S. Patent 3,937,836, discusses the use of specific antibiotic compounds for ruminant feed utilization improvement when given orally to the animal. Specifically, the animal develops rumen function where more propionates in relation to acetates are produced thus

[illegible]

Other references on the use of additives such as monensin have mentioned the need for wise application of these materials because they can be toxic to some animals such as horses. These antibiotics, which are not approved for use in dairy cows, must be administered carefully. In addition, feed intake is initially reduced and monensin can not be added to molasses based supplements which are classic additives to cattle feeds. (Pate, F., "Ionophores Do Not Appear To Work In Molasses Supplements", ONA Reports, November, 1996, 2 pages, Florida Cattleman and Livestock Journal. Lona, R.P. et al, J. Anim. Sci. 75(1)): 2571-2579, 1979.)

Stolle et al, U.S. Patent 4,748,018 is directed to a method of passive immunization of mammals using avian egg yolk antibody against any of a variety of antigens using various methods of administration under various conditions and using various compositions incorporating the antibody, after first developing in the mammal a tolerance for the antibody.

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from eggs and method of production and use thereof for the treatment of infectious or other diseases, and as additives in food for livestock and poultry, cosmetics, and medicines, and in the field of serodiagnosis. Although not explicitly stated, it is apparent that the use of the egg antibody in feeds is to provide an easy means of oral administration of the antibody for the treatment of intestinal infections in livestock or poultry.

Carroll, U.S. Patent 5,196,193 and divisional Patent 5,443,976 are directed to anti-venom compositions containing horse antibody or avian egg yolk antibody for neutralizing snake, spider, scorpion or jelly fish venom.

Lee, U.S. Patent 5,367,054 is directed to methods for large scale purification of egg immunoglobulin for the treatment of infections.

Coleman, U.S. Patent 5,585,098 is directed to a method of oral administration of chicken yolk immunoglobulins to lower somatic cell count in the milk of lactating ruminants.

Stolle et al, U.S. Patent 5,753,268 is directed to an anti-cholesterolemic egg vaccine and method for production and use as a dietary supplement for the treatment of vascular disorders in humans and other animals.

## **SUMMARY OF THE INVENTION**

Broadly stated this invention is directed to a method for the production of a microbial adherence inhibitor for administration to host food animals to substantially prevent the adherence of colony-forming immunogens or haptens in the rumen and /or intestinal tracts of the food animals by first inoculating female birds, in or about to reach their egg laying age, with the particular target immunogen. Then, after a period of time sufficient to permit the production in the bird of antibody to the targeted immunogen, the eggs laid by the birds are harvested. The total antibody-containing contents of the eggs are separated from the shells and dried. The egg

contents may be dried on a feed extender or carrier material. The dried separated egg antibody adherence inhibiting material may be stored or shipped for use when needed.

The target immunogen with which the bird is inoculated depends upon the anticipated use of the inhibitor, a non-disease-causing protein-wasting organism where boosting of feed efficiency is the objective, and a targeted disease-causing organism where the objective is the substantial reduction or elimination of illnesses.

The dried egg contents incorporating the antibody specific to the targeted immunogen is administered to the food animals by distributing the antibody material substantially uniformly throughout an animal feed and then supplying the resulting antibody-containing animal feed to the food animals. When improved feed utilization is the objective, the antibody-containing animal feed is supplied to food animals during the normal finishing schedule prior to slaughter. The substantial prevention of colonization of the targeted organism in the rumen or intestinal tract of the animal will ultimately permit elimination of the organism from the animal. This repression of colonization and elimination of the subject organisms will permit a significant decrease in the wasteful degradation of the dietary protein fed to food production animals. In addition, the resulting decrease in competition to the non-ammonia producing organisms will further enhance the most efficient utilization of feed by the host. (Russell, USDA-ARS, May 1993). When the objective is the elimination of disease-causing organisms from the meat of food animals, the antibody-containing feed is supplied sufficiently before slaughter to substantially prevent adherence of the target immunogen or hapten in the intestinal tract of the animal, and permit elimination of the immunogen or hapten from the animal.

The invention is directed particularly to the production of an adherence inhibitor specific to E. coli 0157:H7 and to the substantial reduction or elimination of gastric illnesses caused by

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this bacterium. The invention is described with particular reference to elimination of illnesses caused by E. coli 0157:H7, but it is to be understood that the invention is not so limited, but is equally applicable to elimination of illnesses caused by other colony-forming immunogens and haptens.

## DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention is based on the concept of specifically inhibiting the ability of colony-forming protein-wasting organisms, such as P. anaerobius, C. sticklandii and C. aminophilum, and colony forming disease-causing organisms, such as E. coli 0157:H7, Listeria, Salmonella and Campylobacter, to adhere in the rumen or intestinal tract of food animals and thus reduce their ability to multiply, grow and colonize. Dietary modifications may be designed to make the rumen and intestinal tract less receptive to the organisms over the lifetime of the animal. While the microbial inhibitor of the present invention may be administered at will by the producer, it is preferred for efficient animal feed utilization that a carefully determined and managed course of administration during the finishing period at the feedlot level be scheduled and followed. Such a predetermined period which takes advantage of the low dose, longer cumulative effect of the inhibitor and which is also easily integratable into current production practices will provide the most economically attractive rate of return through improved animal performance.

For the elimination of disease-causing organisms the inhibitor may be administered either immediately pre-slaughter or over some substantial period of the lifetime of the animal. It is preferred that a carefully determined and managed mid-term period course of administration at the feedlot level be followed. As described, a set pre-slaughter period takes advantage of the low dose, longer cumulative effect, it is easily integratable into current production practices and

is the most economical. It also allows the microorganism to naturally disappear from the mud and manure on the outside of the animal, a significant source of potential contamination at slaughter. Under the current feeding system, food animal feed efficiency is enhanced through the use of ionophores such as monensin, a feed additive marketed under the trade name Rumensin. These are a class of polyester antibiotics approved for feed given to beef cattle and dairy heifers but not approved for use with lactating dairy cows. Most gram-positive organisms are non-specifically vulnerable to the ionophores, antibiotics which can also be quite toxic to the host animal if used improperly. As these antibiotics are not specific, many of the ruminal organisms required to digest the cellulose of ingested plant material may also be affected. The problem with carry over and the development of drug resistant strains of organisms are also major concerns to the industry. The use of broad spectrum antibiotics has further drawbacks including vulnerability to human error, additional cost, consumer resistance and the like. In addition, the monensin type additive can not be administered with commonly used molasses based supplements.

Any organism that colonizes the rumen or alimentary tract of its host must possess the capability of sticking or adhering to that surface in order to multiply and grow. The specific organisms addressed by this invention are no exception to the rule. As other factors such as the need of beneficial organisms for specific enzymes must also be considered, specific reagents are required to reduce the number of targeted organisms in the rumen or intestinal tract while not interfering with other normal flora. The organism inhibitor of this invention strongly interferes with adherence in a highly specific manner and, on a cumulative basis, thereby prevents the targeted organisms from multiplying, growing and colonizing. Through the vehicle of a simple daily feed supplement, the product essentially supplies the host with an antibody preparation

designed not to cure any disease in the animal but to specifically dislodge any resident bacteria in the rumen or alimentary tract and to prevent attachment of any newly introduced numbers of that same bacteria. The microbial inhibitor has no direct effect whatsoever on the ultimate food products and leaves absolutely no undesirable residue in the animal or in the ultimate food products. In addition, since the deleterious organisms are prevented from multiplying; they will over time, for example the 120-day finishing period in the feedlot, disappear through natural degradation from the feedlot environment helping to eliminate that significant potential source of recontamination. The inhibitor product itself can be classified as a natural material of animal origin and as such can be used in almost any kind of feeding program. As the active ingredients are completely natural, they will work well with most feeds and feed additives including molasses based supplements.

All mammals and birds provide similar types of protection which allow for an immediate immune response in their very young offspring until they too acquire the ability to make the antibodies for themselves. More specifically called passive antibody protection, this defense mechanism is passed to the young of mammals through the placenta, the mother's milk or through both. The young of birds, however, receive their passive antibody protection through the store of antibodies placed in the eggs in which they develop from the embryonic stage. Birds, in particular, have the ability to "load up" their eggs as they are formed, with a very large supply of antibodies concentrated many fold over that which is present in the serum of the mother. In addition, avian antibodies are much more stable and resistant to inactivation through digestion than mammalian antibodies, especially under adverse conditions. Once immunized the hen layers the unique IgY types immunoglobulins in the yolk while depositing the common chicken IgM and IgA immunoglobulins in the albumin. The albumin helps

resistance to the whole egg preparations and helps protect the avian antibodies Furthermore, the large quantities of antibodies which are placed in eggs are much more exclusively those specific for the antigens to which the mother has most recently been exposed to and challenged by. This all results in the eggs of birds being a most ideal source for large quantities of economically produced, highly specific and stable antibodies. While the invention is illustrated by the use of chickens to produce avian antibody, other fowl including turkeys, ducks, geese, etc. may be used.

Specifically, groups are obtained of young hen chickens typically Rhode Island Reds, white leghorns, sex-linked hybrid crosses or other breeds suited to large egg size, high volume egg production and ease of handling which are about to reach laying age, about 19 weeks for chickens, on a schedule predetermined by the amount and timing of final product desired resulting in a steady continuous production stream. After a suitable period of isolation and acclimatization of about 2 to 4 weeks, each group will enter into an inoculation program using rehydrated proprietary preparations of specific antigens to which an antibody is desired. The antigens may be obtained from commercial sources such as the American Type Culture Collection (ATCC). The antigen may be injected intra-muscularly, but preferably injected subcutaneously. In approximately four to five weeks, the average egg collected will contain copious amounts of the desired specific antibody in a readily usable and stable form. The chickens may be reinoculated with the targeted antigen throughout the egg laying period to maintain the high antibody level.

Batches of eggs from predetermined groups of chickens are cracked, the contents are separated from the shells and mixed and preferably pasteurized (to eliminate potential pathogenic microorganism from the chicken and thus reduce potential contamination of feed).

The total egg content is dried using standard commercial methods, such as spray drying using ambient or hot air up to 50°C. and tested to determine overall titer or antibody level. The egg contents may be dried alone or on innocuous feed extenders such as dry soy or rice husks or the like. Standard test procedures are used, such as ELISA, or agglutination, or the like. The typical batch is then blended with batches from groups of chickens at other average production levels resulting in a lot of standardized active ingredient. The dried egg antibody microbial inhibitor material may be stored and shipped on carrier materials such as soy bean hulls, boluses and/or tablets. Dependent on the needs and specifications of the feed formulator and the final customer, the final antibody product may include some type of innocuous additive, such as dried whey or dried soy protein powder, dried soy or rice husks or the like for formulation with feed ration. One egg produced and processed by the above procedures will yield a product sufficiently active and stable to provide at least as many as 350 to 700 daily doses of managed protection against specific microbial colonization. This method provides for the first time, an economical, safe and effective means for controlling feed efficiency organisms in beef cattle and dairy herds, and an economical, safe and effective means for controlling E. coli 0157:H7 and other illness-causing organisms in cattle herds.

The present invention specifically addresses feed efficiency as it relates to beef cattle, and by extension dairy cattle and dairy herds, and to the problem of eliminating illness-causing organisms from cattle. However, the concept of preventing microbial adherence has great economic potential for a number of diverse food safety and production application. One such field of application is in feed and water targeting specific undesirable microorganisms. An example of this application would include products to actively inhibit pathogenic or even spoilage microorganisms in animal feed formulated for chickens and other poultry. Another

such field of application is as rinse aid ingredients targeted to specific undesirable microorganisms. Examples of this application include products to actively dislodge pathogenic or even spoilage microorganisms for use in solutions for spot cleaning and rinsing beef carcasses or for chilling poultry after they have been dressed.

The most successful colonizing microorganisms, bacteria, viruses and parasite, etc., have evolved a number of different types of molecules, referred to as “adherins,” on their surfaces which can very tightly stick to one or more types of specific molecules that are part of the host’s various surfaces. The adhesion inhibitor is an avian antibody of extraordinarily high specific activity which can very tightly bind to, coat, cover and obliterate these adherins which attach themselves to their hosts with a lock and key type of fit to very unique chemical structures. In addition to this direct attack, components of the complement system included in most biological fluids, such as blood, lymph, saliva, tears and to some extent intestinal secretions, recognize an antibody attachment as triggers for their many types of defensive activities. Specific antibody attachment and coating combined with the very likely mobilization of many other cellular defense systems, therefore, quickly culminates in the chemical inactivation and ultimately the destruction of the targeted microorganism.

The invention is further illustrated by the following examples:

#### Example 1: Selection of Egg Laying Avian Hens

The strain of egg laying hen may vary with needs and uses. Any egg laying fowl hens may be immunized including chickens, turkeys, ducks, emus or any other fowl. The common strains of egg laying chickens are the preferred and are usually selected for the number of eggs laid per year, size of egg and ease of housing. Rhode Island Red, White Leghorn and Red Sex Linked hybrids are the animals of choice based on egg size (large to ex-large, 50-65 gm) and



were used for the immunization schedules. The ease of handling the animals and the size and uniformity of the eggs along with the number of eggs laid per hen per year were observed. Although any avian egg laying hen could be used, for cost and ease of use these chickens proved to work the best. The Red Sex linked Hybrid gave the most uniformity and greater number of eggs per animal. These animals produce a large to ex-large grade of egg (50-65 gm) and up to 300 eggs a year per hen.

#### Example 2: Preparation of Stock Culture

The American Type Culture Collection E. coli 0157:H7 Stock #43895 was used as the model bacterium. The organism was isolated from raw hamburger and colonizes the cattle. The ATCC Method for rehydration of the stock was followed. The bacterium is rehydrated in 1.0ml of TSB Broth (Tryptase Soy Broth, Becton Dickinson), transferred to 5ml of TSB sterile broth and incubated overnight (approx. 18 hrs) at 37 C. Nice turbid growth was observed. This is used as stock as needed. It was streaked on Sorbitol-MacConkey Agar (Difco) for verification of colony production.

#### Example 3: Preparation of H antigens for Immunogens

The H antigens were selected for development into an immunogen for immunizing the egg laying hens. Certain conditions are used to maintain the optimum growth of the H antigen during culturing to give added concentrations for the prep. Veal Infusion Agar (VIS) and Veal Infusion Broth (VIB, Becton Dickinson) is preferred for H& antigen production. Stock TSB innoculate VIB is incubated at 22-24 C or room temperature for 18 hr. This stimulates flagella development on the bacteria. Flasks layered with VIA are inoculated with VIB culture. Good growth was seen after 22 hours. The product was harvested after 4 days. Flasks are combined by washing off the agar surface with Dulbecco's PBS solution (pH 7.3-7.4). The product is

collected in tubes. Density is checked using spectrophotometric enumeration and McFarland nephelometer standards. Approximately  $3 \times 10^{12}$ /ml in Stock. Motility is checked with Motility agar slant (Northeast Laboratory Services). Stock is diluted to concentration of approx.  $1 \times 10^9$  per ml in PBS and stirred for 1 hr at room temperature. The Flagella is removed from the outside of the bacteria. Supernatant is collected using centrifugation. Pellet of whole bacteria is separated from the supernatant. Dry weight approx. 14.7mg/ml is determined and the material is used as Stock Immunogen for H antigen. It is diluted to 1 mg/ml in PBS and heated for 30 minutes at 60-70 C. This helps keep contamination down to a minimum. Thioglycollate broth is inoculated to check for growth and animals are inoculated with immunogen.

#### Example 4: Preparation of O Antigen for Immunogens

Brain Heart Infusion (BHI, acumedia) is used to stimulate the O antigens on the bacterium. Stock TSB inoculate BHI Broth is formed and incubated at 37°C for 18 hrs. This stimulates somatic antigen development on the bacteria. Flasks containing BHI Broth are inoculated with BHI Broth culture. While stirring slowly, flasks are incubated at 37°C. Good growth is seen after 22 hours. Flasks are combined and the material is harvested using centrifugation and sterile saline (0.9%) at approx. 3000rpm for 30 minutes. The harvest is collected in tubes. Density is checked using spectrophotometric enumeration and McFarland nephelometer standards. The material is diluted to approximately  $1 \times 10^9$  per ml. 4% sodium deoxycholate (Difco) solution is added as a 1:1 ratio with culture in 0.9% sterile saline (Herzberg, 1972) and stirred for approx. 18 hrs at room temperature (22-24C). The material is centrifuged to remove whole cells. Supernatant is used as stock for O antigen. Dry weight is determined at approximately 14.9mg/ml. the product is diluted in sterile PBS, pH 7.4 to 1mg/ml for O Immunogen.

#### Example 5: Preparation of WC Antigen for Immunogens

Tryptic Soy Broth (TSB, Northeast Laboratory Services) plus Yeast Extract (BBL) is used for Whole Cell (WC) Antigen Production. TSB plus Yeast Extract 0.6% Broth is inoculated from TSB Stock and incubated at 37°C for 18 hrs. This stimulates somatic and other surface antigens to development on the bacteria. Flasks are inoculated with TSB with Yeast Extract Broth. While stirring slowly, it is incubated at 37°C. Good growth is seen after 22 hours. The flasks are combined and the product is harvested using centrifugation at approx. 3000 rpm for 30 minutes and collected in tubes. The product is resuspended in sterile PBS, pH 7.4. Density is checked using spectrophotometric enumeration and McFarland nephelometer standards. Dry weight is approximately 19.7mg/ml. The product is diluted to approximately  $2 \times 10^9$  per ml or 2mg/ml dry weight, and 0.6% formaldehyde solution in PBS is added as a 1:1 ratio with culture and stirred for approx. 18 hrs at RT (22-24C) to fix cells. Thioglycollate broth is inoculated to check for growth and pH of preparation (pH 7-7.4) is checked. the supernatant is used as for WC antigen. The stock is diluted in PBS, pH 7.4 to 1mg/ml for WC immunogen.

#### Example 6: Preparation of A antigen for Immunogen

The Minca Medium is used for A Antigen Production. It is a standard medium for stimulating the pili and related adherin antigens. Stock TSB Minca Medium Broth (Inf. Immun., Feb. 1977, 676-678) is inoculated and incubated at 37°C for 18 hrs. This stimulates adhesion antigen development on the bacteria. Flasks are inoculated with Minca Medium Broth and while stirring slowly is incubated at 37°C. Good growth is seen after 18 hours. The flasks are combined and the product is harvested using centrifugation at approx. 2500 rpm for 30 minutes and collected in tubes. The pellet is resuspended in PBS and stirred with a stir bar for 1

hour at 22-24°C (RT). This removes the flagella. The product is collected in tubes and the pellet is resuspended in PBS and 0.01% Tween 20™, transferred to Waring Blender in cold (4°C) at low speed for 30 minutes. Density is checked using spectrophotometric enumeration and McFarland nephelometer standards. The product is centrifuged to remove whole cells. the supernatant is used as stock for A antigen. It may be heated at 60°C for 40 min. to inactivate if needed. Gentamycin is added at 50 µg/ml as preservative. Thioglycollate broth is inoculated to check for growth. Dry weight is determined at approximately 10.6 mg/ml. The product is diluted with PBS, pH 7.4 to 1 mg/ml for A Immunogen.

#### Example 7: Preparation of P antigen for Immunogen

The Reinforced Clostridial Medium is used for P antigen Production. It is a standard medium for stimulating adherence antigens for *Peptostreptococcus anaerobius*. These cultures must be grown under strict anaerobic conditions. The stock culture is grown according to ATCC for #49031. As with other organisms, subcultures and grown in small amounts. Thioglycollate Media (Difco) is inoculated with the stock and incubated for 48 hrs. Flasks are inoculated with Reinforced Clostridial Medium Broth. The medium is covered with a mixture of anaerobic gas. Flasks are combined and the product is harvested using centrifugation at approximately 2500 rpm for 30 minutes, collected in tubes and run at low speed for 30 minutes. Density is checked. The product is centrifuged to remove whole cells. the supernatant is used as stock for P antigen. It is heated at 60°C for 40 min. to inactivate if needed. Dry weight is determined. Approximately 20.5 mg/ml. The product is diluted with PBS, pH 7.4 to 1 mg/ml for P Immunogen.

#### Example 8: Preparation of CS antigen for Immunogen

The Reinforced Clostridial Medium is used for CS Antigen Production. It is a standard medium for stimulating adherence antigens for *Clostridium sticklandii*. These cultures must be

grown under strict anaerobic conditions. The stock culture is grown according to ATCC for #12662. As with other organisms, subcultures are grown in small amounts. Thioglycollate Media (Difco) is inoculated with the stock and incubated for 48 hrs. Flasks are inoculated with Reinforced Clostridial Medium Broth. The medium is covered with a mixture of anaerobic gas. Flasks are combined and the product is harvested using centrifugation at approximately 2500 rpm for 30 minutes. The product is collected in tubes and spun at low speed for 30 minutes. Density is checked using spectrophotometric enumeration and McFarland nephelometer standards. The product is centrifuged to remove whole cells. The supernatant is used as stock for CS antigen. It is heated at 60°C for 40 min. to inactivate if needed. Dry weight is determined at approximately 22mg/ml. The product is diluted with PBS, pH 7.4 to 1mg/ml for CS Immunogen.

#### Example 9: Preparation of CA antigen for Immunogen

The Reinforced Clostridial Medium is used for CA Antigen Production. It is a standard medium for stimulating adherence antigens for *Clostridium aminophilus*. These cultures must be grown under strict anaerobic conditions. The stock culture is grown according to ATCC for #49906. As with other organisms, subcultures are grown in small amounts. Thioglycollate Media (Difco) is inoculated with the stock and incubated for 48 hrs. Flasks are inoculated with Reinforced Clostridial Medium Broth. The medium is covered with a mixture of anaerobic gas. Flasks are combined and the product is harvested using centrifugation at approx 2500 rpm for 30 minutes. The product is collected in tubes and spun at low speed for 30 minutes. Density is checked using spectrophotometer enumeration and McFarland nephelometer standards. The product is centrifuged to remove whole cells. The supernatant is used as stock for CA antigen. It is heated at 60°C for 40 min. to inactivate if needed. Dry weight is determined at

approximately 20.5mg/ml. The product is diluted with PBS, pH 7.4 to 1mg/ml for CA

Immunogen.

Example 10: Preparation of ELISA Plates using H, O, WC and A Antigens for monitoring antibodies in eggs, chickens and feed.

H, O, WC and A ELISA: Ninety six well assay plate (flat bottom Costar®) were coated using 100µl/ml with various concentration of antigens (H, A, O, or WC or combination: 10µg-200µg/ml) in carbonate buffer, pH 9.6. Plates were incubated between 22-37°C for up to 18 hrs. The wells were aspirated to prevent cross-contamination. The plates were blocked with 390µl/well of 0.5% BSA and incubated at 37°C for 1 hr. Plates were coated using alternative rows of positive or negative for controls. Plates are rinsed 1X with wash buffer containing Tween™20. One hundred microliters per well of diluted sample are added to wells in duplicate wells. Incubated at 37°C for one hour. Goat anti-Chicken IgG conjugate with Horseradish peroxidase (Kirkegard and Perry laboratories; 1:1000-1:3000) was added. After 1 hr incubation, the substrate (TMB, KPL) was added according to manufacturer's instructions and the reaction is stopped after 10 minutes with 0.1M phosphoric acid. Optical densities of the wells were determined in Dynatech ELISA Reader at 450nm and the information was recorded for further data analysis.

Example 11: Analysis of Individual Eggs and Serum Over Time

Eggs were selected at various periods in the immunization period for monitoring antibody responses to the specific antigens. Selected chickens were monitored at day 0 and continued on a monthly basis after the 4<sup>th</sup> month. The whole egg was collected from the shell and then a 1 ml sample was taken. This sample was then extracted with buffer to analysis the antibody content. The standard ELISAs for the H, O, WC and A Immunogens were used for analyse. The negative readings were subtracted from the OD readings. Serum samples were collected from each animal two weeks after the fourth immunogen injection.

The data given in the table below are examples of the results obtained over the first four months.

Egg Sample Date	H Chicken	O Chicken	WC Chicken	A Chicken
1 day:After first injection	0.03OD	Neg	0.05OD	Neg
1 month	0.60OD	Neg	0.05OD	Neg
5 weeks	0.74	ND	ND	ND
2 months	1.22OD	1.11OD	0.88OD	0.79OD
3 months	1.00OD	1.4OD	0.99OD	1.4OD
4 months	1.16OD	1.4OD	0.94OD	1.22OD
Serum: 1 month	1.4OD	0.91OD	1.17OD	0.97OD

Example 12: Preparation of ELISA Plates using P, CS and CA Antigens for Monitoring antibodies in eggs, chickens and feed.

P, CS and CA ELISA: Ninety six well assay plate (flat bottom Costar®) were coated using 100µl/ml with various concentrations of antigens (P, CS, CA or combination:10µg-200µg/ml) in carbonate buffer, pH 9.6. Plates were incubated between 22-37°C for up to 18 hrs. The wells were aspirated to prevent cross-contamination. The plates were blocked with 390µl/well of 0.5% BSA and incubated at 37° C for 1 hr. Plates were coated using alternative rows of positive or negative for controls. Plates are rinsed 1X with wash buffer containing Tween™20. One hundred microliters per well of diluted sample are added to wells in duplicate wells, and incubated at 37° C for one hour. Goat anti-Chicken IgG conjugate with Horseradish peroxidase (Kirkegard and Perry laboratories; 1:1000-1:3000) was added. After 1hr incubation, the substrate (TMB, KPL) was added according to manufacturer's instructions and the reaction is stopped after 10 minutes with 0.1M phosphoric acid. Optical densities of the wells were determined in Dynatech ELISA Reader at 450nm and the information was recorded for further data analysis.

Example 13: Immunization of Chicken with H Immunogen:

Six selected egg laying hens, 3 White Leghorn and 3 Rhode Island Red approximately 19 weeks old were injected with the stock H Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, 4 out of the 6 hens produced excellent antibodies in the eggs. ELISA H readings averaged 1.00 OD for 1:10,000 dilution and 0.265OD for 1:50,000. Leghorn hen's did not do as well but all 3 Rhode Island did well. After six weeks the average ELISA H reading was 1.40 OD for 1:20,000 dilution with all chickens responding.

Example 14: Immunization of Chicken with O Immunogen:

Six selected egg laying hens, 6 White Leghorn, approximately 19 weeks old were injected with the stock O Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, 5 out of the 6 hens produced excellent antibodies in the eggs. ELISA O readings averaged 1.42 OD for 1:10,000 dilution and 0.680OD for 1:50,000. Leghorn hens did not do as well but all 3 Rhode Island did well. After six weeks the average ELISA O reading was 1.15 OD for 1:20,000 dilution with still 5 chickens responding.

Example 15: Immunization of Chicken with WC Immunogen:

Six selected egg laying hens, 6 Rhode Island Red, approximately 19 weeks old were injected with the stock WC Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last



### Example 13: Immunization of Chicken with H Immunogen:

Six selected egg laying hens, 3 White Leghorn and 3 Rhode Island Red approximately 19 weeks old were injected with the stock H Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, 4 out of the 6 hens produced excellent antibodies in the eggs. ELISA H readings averaged 1.00 OD for 1:10,000 dilution and 0.265OD for 1:50,000. Leghorn hen's did not do as well but all 3 Rhode Island did well. After six weeks the average ELISA H reading was 1.40 OD for 1:20,000 dilution with all chickens responding.

### Example 14: Immunization of Chicken with O Immunogen:

Six selected egg laying hens, 6 White Leghorn, approximately 19 weeks old were injected with the stock O Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, 5 out of the 6 hens produced excellent antibodies in the eggs. ELISA O readings averaged 1.42 OD for 1:10,000 dilution and 0.680OD for 1:50,000. Leghorn hens did not do as well but all 3 Rhode Island did well. After six weeks the average ELISA O reading was 1.15 OD for 1:20,000 dilution with still 5 chickens responding.

### Example 15: Immunization of Chicken with WC Immunogen:

Six selected egg laying hens, 6 Rhode Island Red, approximately 19 weeks old were injected with the stock WC Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last

initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, 4 out of the 6 hens produced excellent antibodies in the eggs. ELISA WC readings averaged 0.95 OD for 1:10,000 dilution and 0.250OD for 1:50,000. After six weeks the average ELISA O reading was 0.95 OD for 1:20,000 dilution with still 5 chickens responding.

#### Example 16: Immunization of Chicken with A Immunogen:

Six selected egg laying hens, 6 White Leghorn, approximately 19 weeks old were injected with the stock A Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug were given in each booster (every 6 months). Within four weeks, 5 out of the 6 hens produced excellent antibodies in the eggs. ELISA A readings averaged 1.40 OD for 1:10,000 dilution and 0.576 OD for 1:50,000. After six weeks the average ELISA A reading was 1.15 OD for 1:20,000 dilution with still all chickens responding.

#### Example 17: Immunization of Chicken with P Immunogen:

Six selected egg laying hens, White Leghorn , approximately 19 weeks old were injected with the stock P Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug were given in each booster (every 6 months). Within four weeks, 5 out of the 6 hens produced excellent antibodies in the eggs.

#### Example 18: Immunization of Chicken with CS Immunogen:

Six selected egg laying hens, White Leghorn, approximately 19 weeks old were injected with the stock CS Immunogen. Four injections (500ug, 100ug, 200ug, and

250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, All 5 out of 6 hens produced excellent antibodies in the eggs.

#### Example 19: Immunization of Chicken with CA Immunogen:

Six selected egg laying hens, Red Sex-Linked Hybrids, approximately 19 weeks old were injected with the stock CA Immunogen. Four injections (500ug, 100ug, 200ug, and 250ug) were given 1 week apart. A serum sample was collected two weeks after the last initial injection. If boosters were needed, 100ug was given in each booster (every 6 months). Within four weeks, All 6 hens produced excellent antibodies in the eggs.

#### Example 20: Preparation of Stock Production Whole Egg reagents

Selected hens were combined from all four immunogen groups for E. coli O157:H7 or three immunogen groups for the anaerobes, to be used to produce production batches of whole egg reagents. Sterling U.S. Patent 5,753,228 presents an excellent review of uses for the selection of eggs and storage of same. The eggs were randomized and shell removed. The whole egg is mixed well and pasteurized using standard conditions (60°C (140°F) for 3.5min), Charley, H. and C. Weaver, 3<sup>rd</sup> edition, Foods: a scientific approach, Merrill-Prentice Hall, p350, 1998). Once pasteurized, samples were tested for activity and stored at 4° C until dried or sprayed onto carriers. Samples of 250µl were analyzed.

Examples of results for ELISAs are given:

Pasteurized Whole Egg: E. coli O157:H7

Immunogen	Dilution	O.D.
WC	500	0.532
WC	2500	0.113
H	500	0.466
H	2500	0.115
O	500	0.338
O	2500	0.128
A	500	0.588
A	2500	0.155

Pasteurized Whole Egg: Anaerobes

Immunogen	Dilution	Batch #1	Batch#2	Batch #3
CA	100	0.339	0.275	0.627
CA	500	0.104	0.296	0.201
P	100	0.724	0.882	0.576
P	500	0.248	0.594	0.651
CS	100	0.457	0.268	0.650
CS	500	0.304	0.143	0.476

Example 21: Coating of Feed Additive Carriers

Although whole egg can be dispensed in water supplies, or in a dried format as whole powdered egg, use of a carrier helps distribute the material in a uniform method. This makes it easier for mixing with standard feeds. A number of carriers can be used to provide a vehicle as a feed additive as needed. Soy hulls in crude, refined and pelted format, rice hulls, corn, cottonseed hulls, distilled dried grains, beet pulp or any other. The production pasteurized whole egg prep is coated on to the carrier and either fed directly to the animals or dried to 10-15% moisture. Approximately 1000ml of whole, pasteurized egg is sprayed on 50lbs of pelleted soybean hulls. The preferred carrier for cattle is pelleted soybean hulls while for young swine the fines from the pelleted soybean hulls. The feed additive is mixed with the standard animal feed. The preferred level is 10-15lbs of feed additive to 2000lbs of animal feed.

### Example 22: Analysis of Feed Additive samples after coating with reagents

Samples were collected from batches of Feed Additive after they were coated on to the carriers. The samples were analyzed and the results are as follows:

Product Name	Moisture %	Protein %	Fat %	Fiber, crude %
Crude Soybean Hulls, uncoated	11.59	26.76	9.10	18.63
CAMAS EYE 0157 Crude soybean Hulls	12.35	25.67	8.26	19.46
CAMAS EYE * Control Crude Soybean hulls	12.06	24.89	9.92	20.38
Soybean Pellets uncoated	11.65	9.89	2.43	33.47
CAMAS EYE Efficiency Pellets	12.37	10.19	2.57	33.12

\* CAMAS EYE identifies inhibitors produced according to the present invention

### Example 23: Analysis of Production Eggs over Time: E. coli O157:H7

Samples of the Whole Egg preparations were analyzed using the ELISA systems for H, O, WC and A immunogens to monitor activity over time after the initial immunization schedule was completed. Selected animals from each group were placed into the Production group. The average ELISA OD readings (Negative subtracted) for the fourth through the six months are given in the table below. The eggs were sampled using 250μ of the whole egg and diluted 1:500 and 1:2500 in PBS buffer and then run in the appropriate ELISA to determine the average OD reading at each dilution. The negative control readings are subtracted from each reading. The immunogens showed different responses in the animals along with good specificity. The A immunogen gave the best responses in these tests. Data for these immunogens over time is given below:

Immuogen	Fourth Month	Fifth Month	Six Month
H : 1:500	0.388	0.848	0.718
1:2500	0.085	0.237	0.195
O: 1:500	0.593	0.792	0.704
1:2500	0.147	0.294	0.184
WC: 1:500	0.398	0.730	0.578
1:2500	0.062	0.273	0.130
A: 1:500	0.700	1.014	0.909
1:2500	0.102	0.305	0.224

#### Example 24: Analysis of Production Eggs over Time: Feed Efficiency

Samples of the Whole Egg preparations were analyzed using the ELISA systems for P, CS, and CA immunogens to monitor activity over time after the initial immunization schedule was completed. Selected animals from each group where placed into the Production group. The average ELISA OD readings for the fourth through the six months are given in the table below. The eggs were sampled using 250µ of the whole egg and diluted 1:500 and 1:2500 in PBS buffer and then run in the appropriate ELISA to determine the average OD reading at each dilution. The negative control readings are subtracted from each reading. The immunogens showed different responses in the animals along with good specificity.

Immuogen	Fourth Month	Fifth Month	Six Month
P : 1:500	1.182OD	1.128OD	0.942OD
1:2500	0.785OD	0.489OD	0.343OD
CS: 1:500	0.843OD	0.989OD	0.582OD
1:2500	0.318OD	0.356OD	0.187OD
CA: 1:500	1.156OD	1.087OD	0.998OD
1:2500	0409OD	0.282OD	0.507OD

#### Example 25: Analysis of Feed Additives for Antibody Activity: *E. coli* O157:H7

Samples of the coated hulls were analyzed using the ELISA systems for H, O, WC and A immunogens to monitor activity after pasteurizing, spraying, drying and storage.

Good antibody response was recorded after the processing of the Production Whole Egg batches and drying on crude soybean hulls. Data for two batches is given below:

Batch: Coated Hulls	WC Immunogen	H Immuogen	O Immunogen	A immunogen
Batch #1 1:10	0.673 OD	1.103 OD	1.105 OD	1.299 OD
1:100	0.106 OD	0.236 OD	0.229 OD	0.302 OD
Batch #2 1:10	1.174 OD	1.291 OD	1.180 OD	1.224 OD
1:100	0.177 OD	0.396 OD	0.327 OD	0.458 OD

#### Example 26: Analysis of Feed Additives for Antibody Activity: Feed Efficiency

Samples of the coated hulls were analyzed using the ELISA systems for P,CS, and CA immunogens to monitor activity after pasteurizing, spraying, drying and storage.

Good antibody response was recorded after the processing of the Production Whole Egg batches and drying on crude soybean hulls. One gram samples of the 15lbs of coated hulls was extracted and analyzed. Data for three batches is given in the table below:

Batch: Coated Hulls	P Immunogen	CS Immuogen	CA Immunogen
Batch #1 1:100	0.067OD	0.289OD	0.051OD
1:500	0.057OD	0.131OD	0.037OD
Batch #2 1:100	0.028OD	0.039OD	0.095OD
1:500	0.049OD	0.015OD	0.021OD
Batch#3 1:100	0.046OD	0.115OD	0.136OD
1:500	0.012OD	0.055OD	0.012OD

#### Example 27: Recovery of Active Antibody and Egg Protein After Feed Mix

Bags of coated soybean refined hulls were coated with the production whole egg reagent containing anti-E. coli O157:H7 adherence inhibitors. One bag of feed additive (15lbs) was added to 2000lbs of Standard Cattle Feed. Control Feed Additive was produced with whole eggs from free ranging chickens. Soybean hulls were coated with this preparation and mixed as the Test Feed Additive containing the specific antibodies. Samples of the mixed Feed were collected and analyzed for active antibody to the ELISA

WC immunogen as well as a commercial ELISA for detecting Egg Protein in food (Veratox® Quantitative Egg Allergen Test, Neogen). The data is given in the chart below for two batches of feed ration.

Mixed Feed	First Batch	Second Batch
Test Feed-Additive: 1:6000	0.172 OD 0.009OD	0.112OD 0.036
1:12000		
Control Feed-No Additive	0.049 0.005	Neg. Neg.
1:6000		
1:12000		
Test Feed-Additive: Egg Protein	0.958 OD 17ppm	1.268OD >20ppm
Control Feed-No Additive: Egg Protein	0.800OD 15 ppm	1.050OD 20ppm

#### Example 28: Feeding of Cattle

Two groups of cattle were feed either the O157:H7 feed additive (coated onto refined soybean hulls) or control feed additive (coated with control eggs and no specific adherence inhibitors). The animals were feed at a rate of 15lbs of feed additive per 2000 lbs feed. They average 10lbs per animal per day. Animals weighed approximately 1000lbs when they started and over 1400 when sent to market. All animals looked very healthy with the Test animals eating more feed during the 87days. Five of the test animals were positive during the start of the experiment for E. coli O157:H7 and only one of the control animals. Within 30 days on feed additive all test animals were negative for E. coli O157:H7 and stayed negative for three consecutive samples over a 30 day period. Standard protocols were followed for sampling. All animals were ear-tagged and placed



in separate pens. Animals were sampled on a weekly basis for the first month and then bi-weekly after that until they were shipped to market. Grab samples were taken from the rectum and placed into sterile labeled bags. All samples were held on ice until processed in the lab. All samples were processed within 4 hours of collection each day. The fecal samples were diluted with TSB with 0.6% yeast extract. Dilutions of the mixture were streaked onto Sorbitol-MacConkey's agar with or without cefixime-tellurite supplement (Dynal®). Colorless colonies are picked for further testing. A latex agglutination test was used to identify E. coli serogroup O157 (Oxoid dry Spot™ E. coli 0157). If positive, then individual colonies were selected for further isolation on SMC agar streak plates. Isolated colonies were run on the commercial EIA for EH E. coli O157 (Binax, NOW® EH E. coli O157). Biochemical confirmation can be done with API-20E (Analytab Products). (Appl. Environ. Microbiol. 62(7)2567-2570, 1996, J. Clin. Micro. 36(10):3112, 1998)

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One of the most startling and distressing characteristics of E. coli 0157:H7 is the small number of microorganisms necessary to produce cases of human illness. By way of example, at least 10,000 of the more virulent Salmonella serotypes but as few as ten E. coli 0157:H7 are required to cause a person to become symptomatic. Therefore, one animal hosting or externally contaminated with the microorganism can, when slaughtered, affect as much as 16 tons of ground beef to the extent that a single helping of the product could result in illness if improperly prepared. Although the probability of any one animal hosting the microorganism at any one time is low, the probability of its presence in any one particular feedlot is high.

There are presently three different methods for protecting the consumer from the E. coli 0157:H7 threat which have been officially recognized. The three methods are (1) thorough cooking, (2) steam pasteurization and (3) irradiation, all of which have specific drawbacks, including human and mechanical error, cost, consumer resistance, and the like.

Any microorganism which colonizes the alimentary tract of its host must possess the capability of sticking or adhering to that surface in order to multiply. E coli 0157:H7 is no exception to this rule. The adherence inhibitor of this invention strongly interferes with adherence and, on a cumulative basis, thereby prevents the specific targeted microorganism from colonizing and multiplying. Through the vehicle of a simple daily feed additive, the product essentially supplies the host with a specific antibody preparation designed not to cure any

disease in the animal (cattle are essentially unaffected by E. coli 0157:H7 being only transitory hosts) but merely to dislodge any resident bacteria and to prevent the attachment of any newly introduced bacteria in the alimentary tract. The adherence inhibitor has no direct effect on the host itself, leaves absolutely no undesirable residue in the animals and thus has no effect whatsoever on the ultimate food products. In addition, since the microorganism is prevented from multiplying, it will over time (for example the 120 day finishing period in the feedlot) disappear through natural degradation from the mud and manure coating the animal, eliminating this significant potential source of contamination at slaughter. Properly managed, the risk of cross contaminating other food sources through feedlot runoff or by the application of manure as fertilizer is also essentially eliminated.

It is apparent that many modifications and variations of this invention as hereinbefore set forth may be made without departing from the spirit and scope thereof. The specific embodiments described are given by way of example only and the invention is limited only by the terms of the appended claims.

## CLAIMS

1. A method for the production of a microbial adherence inhibitor for administration to food animals to substantially prevent the adherence of targeted colony-forming immunogens in the rumen or intestinal tracts of said food animals, which method comprises:

A. Inoculating female birds, in or about to reach their egg laying age, with the particular target colony-forming immunogen.

B. After a period of time sufficient to permit the production in the bird of antibody to the targeted immunogen, harvesting the eggs laid by the birds;

C. Separating the antibody-containing contents of said eggs from the shells; and

D. Drying said separated egg antibody adherence inhibiting material.

2. A method according to claim 1 wherein said colony-forming immunogen is one known to decrease an animal's ability to utilize dietary protein.

3. A method according to claim 1 wherein said colony-forming immunogen is one known to cause food borne illness in humans.

4. A microbial adherence inhibitor produced by the method of claim 1.

5. A microbial adherence inhibitor produced by the method of claim 2.

6. A microbial adherence inhibitor produced by the method of claim 3.

7. A microbial adherence inhibitor for administration to food animals substantially preventing the adherence of targeted colony-forming immunogens in the rumen or intestinal tracts of said animals comprising dried egg contents incorporating antibody specific to said targeted immunogen.

8. A microbial adherence inhibitor according to claim 7 wherein said colony-forming immunogen is one known to decrease an animal's ability to utilize dietary protein.

9. A microbial adherence inhibitor according to claim 1 wherein said colony-forming immunogen is one known to cause food borne illness in humans.

10. A method of promoting the growth of food animals by decreasing the waste of dietary protein caused by the presence of colony-forming protein-wasting immunogens in the rumen or intestinal tracts of food animals by inhibiting the ability of the immunogen to adhere to the rumen or intestinal tracts of animals to reduce the ability of the immunogen to multiply, said method comprising:

A. Inoculating female birds, in or about to reach their egg laying age, with the particular targeted protein-wasting immunogen;

B. After a period of time sufficient to permit the production in the bird of antibody to the targeted immunogen, harvesting the eggs laid by the birds;

C. Separating the antibody-containing contents of said eggs from the shells;

D. Drying said separated egg antibody material;

E. Distributing the resulting dried egg antibody product substantially uniformly through an animal feed or water; and

F. Supplying the resulting antibody-containing animal feed or water to food animals to substantially prevent adherence of the targeted immunogen in the intestinal tract of the animal.

11. A method according to claim 10 wherein said protein-wasting immunogen is selected from the class consisting of *P. anaerobius*, *C. sticklandii* and *C. aminophilum*.

12. A method for substantially reducing or eliminating the incidence of illnesses caused by the presence of targeted colony-forming illness-causing immunogens in meat by inhibiting the ability of the immunogen to adhere to the rumen or intestinal tracts of food animals to reduce the ability of the immunogen to multiply, said method comprising:

A. Inoculating female birds, in or about to reach their egg laying age, with the particular targeted illness-causing immunogen;

B. After a period of time sufficient to permit the production in the bird of antibody to the targeted immunogen, harvesting the eggs laid by the birds;

- C. Separating the antibody-containing contents of said eggs from the shells;
- D. Drying said separated egg antibody material;
- E. Distributing the resulting dried egg antibody product substantially uniformly through an animal feed or water; and
- F. Supplying the resulting antibody-containing animal feed or water to food animals to substantially prevent adherence of the targeted immunogen in the intestinal tract of the animal.

13. A method according to claim 12 wherein said illness-causing immunogen is selected from the class consisting of E. coli, Listeria, Salmonella and Campylobacter.

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# DECLARATION FOR PATENT APPLICATION

Docket No. C150.12.3B

As a below named inventors I (we) hereby declare that:

My residence, post office address and citizenship are as stated below next to my (our) names.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled IMMUNOGEN ADHERENCE INHIBITOR AND METHOD OF MAKING AND USING SAME the specification of which:

(check one) X is attached hereto.  
       was filed on        as  
 Application Serial No.         
 and was amended on        (if applicable).

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

Provisional	Filing Date	Status
Provisional 60/143,985	July 15, 1999	Pending
(Application Serial No.)	(Filing Date)	Status-patented, pending, abandoned)
Provisional 60/201,268	May 2, 2000	Pending
(Application Serial No.)	(Filing Date)	Status-patented, pending, abandoned)

I hereby appoint the following attorney(s) to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith:

L. Paul Burd - Registration Number 17,048  
 Richard O. Bartz - Registration Number 20,468  
 Robert W. Gutenkauf - Registration Number 25,681

Address all telephone calls to L. PAUL BURD at telephone no. (612) 332-6581 or fax at (612) 332-6584. All correspondence to 1300 Foshay Tower, Minneapolis, Minnesota 55402.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Full name of sole or first inventor PETER (nmi) NASH  
 Inventor's signature Peter Nash Date JULY 13, 2000  
 Residence EDEN PRAIRIE, MINNESOTA Citizenship UNITED STATES  
 Post Office Address 1313 Fifth Street SE, Suite 219, Minneapolis, MN 55414  
 Full name of joint or second inventor JOHN W. ROSEVEAR, deceased, by his legal representative  
 Inventor's signature Donald L. Robinson Date JULY 13, 2000  
 Residence EDINA, MINNESOTA Citizenship UNITED STATES  
 Post Office Address 1313 Fifth Street SE, Suite 219, Minneapolis, MN 55414  
 Full name of joint or third inventor DONALD L. ROBINSON  
 Inventor's signature Donald L. Robinson Date JULY 13, 2000  
 Residence BLOOMINGTON, MINNESOTA Citizenship UNITED STATES  
 Post Office Address 1313 Fifth Street SE, Suite 219, Minneapolis, MN 55414